

A plant flavonol and genetic suppressors rescue a pathogenic mutation associated with kinesin in neurons

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KIF1A, a microtubule-based motor protein responsible for axonal transport, is linked to a group of neurological disorders known as KIF1A-associated neurological disorder (KAND). Current therapeutic options for KAND are limited. Here, we introduced the clinically relevant KIF1A(R11Q) variant into the *Caenorhabditis elegans* homolog UNC-104, resulting in uncoordinated animal behaviors. Through genetic suppressor screens, we identified intragenic mutations in UNC-104's motor domain that rescued synaptic vesicle localization and coordinated movement. We showed that two suppressor mutations partially recovered motor activity in vitro by counteracting the structural defect caused by R11Q at KIF1A's nucleotide-binding pocket. We found that supplementation with fisetin, a plant flavonol, improved KIF1A(R11Q) worms' movement and morphology. Notably, our biochemical and single-molecule assays revealed that fisetin directly restored the ATPase activity and processive movement of human KIF1A(R11Q) without affecting wild-type KIF1A. These findings suggest fisetin as a potential intervention for enhancing KIF1A(R11Q) activity and alleviating associated defects in KAND.

kinesin | axonal transport | neurological disorders | C. elegans

Axonal transport, facilitated by microtubules (MTs), is essential for neural development and function (1–3). It involves bidirectional movement mediated by kinesin superfamily proteins (KIFs) and cytoplasmic dynein-1 (4–6). Kinesin-3 family motors, such as UNC-104, transport synaptic vesicles (SVs) from the cell body to the axon (7, 8). Mutations in UNC-104 result in abnormal SV localization and accumulation (9, 10). UNC-104/KIF1A consists of various domains, including a motor domain, neck coil domain (NC), coiled-coil domains, forkhead-associated (FHA) domains, and a C-terminal pleckstrin homology (PH) domain (11–13). Genetic links have been established between KIF1A mutations and impaired axonal transport, leading to neurodegeneration and KIF1A-associated neurological disorders (KAND) (14–19). Numerous KIF1A mutations have been identified in patients, associated with conditions ranging from congenital neuropathies to hereditary spastic paraplegia (SPG) (16–18, 20–23). However, effective interventions for these mutations and KAND therapies are still largely unknown.

Caenorhabditis elegans has emerged as a valuable genetic model for studying KAND. The functional defects observed in the *unc-104* mutant were rescued by introducing the human homolog KIF1A into *C. elegans* (24), indicating the evolutionary conservation of these motor proteins. Recent research has utilized genome editing techniques to introduce KAND-associated mutations into the *C. elegans* genome, allowing for the systematic analysis of axonal transport and animal behavior (25, 26). Complemented by biochemical and single-molecule assays conducted on KIF1A in vitro, these studies shed light on the dominant nature of de novo KAND variants and reveal that certain mutations lead to an abnormal increase in KIF1A motor activity (25, 26). This suggests that hyperactivation of the motor, rather than loss-of-function, may contribute to motor neuron diseases.

C. elegans serves as an excellent platform not only for modeling KAND mutations but also for conducting genetic suppressor screens, offering valuable insights into potential interventions for KAND. For instance, one screen uncovered UNC-16, a kinesin cargo adaptor protein, and specific amino acid substitutions in the motor stalk region, as intergenic suppressors capable of restoring the phenotype of an *unc-104* gene mutant allele, designated *e1265*. The *unc-104 (e1265)* mutant animal possesses a D1497N mutation in the motor's PH domain, leading to aberrant retention of synaptic vesicles within the cell body (27, 28). Another unbiased screen identified a gain-of-function allele of *unc-104* as a suppressor that restored presynaptic patterning and axonal transport disrupted by the loss of ARL-8, an Arf-like small G protein (29). These findings demonstrate the feasibility of isolating suppressors for *unc-104* from both loss- and gain-of-function contexts. Additionally, a recent screen successfully

Significance

This study offers important insights into KIF1A-associated neurological disorder (KAND), a group of neurological disorders connected to KIF1A, a microtubule-based motor protein responsible for axonal transport. With limited therapeutic options for KAND, alternative interventions are crucial. Introducing the KIF1A(R11Q) variant into Caenorhabditis elegans reproduced KAND symptoms of impaired coordination and movement. Genetic suppressor screens identified intragenic mutations in UNC-104 that restored synaptic vesicle localization and coordinated movement. Fisetin, a plant flavonol, improved movement and morphology in KIF1A(R11Q) worms. Biochemical assays demonstrated fisetin's ability to directly restore impaired KIF1A(R11Q) activity without affecting wild-type KIF1A. These findings highlight fisetin as a potential therapeutic intervention for enhancing KIF1A(R11Q) activity and alleviating associated defects in KAND.

The authors declare no competing interest.

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identified a mutation that partially restored the motility of a clinical KIF1A variant with reduced motor activity (26).

This study investigates the impact of the R11Q mutation in the KIF1A motor domain, which has been observed in a patient with combined autism spectrum disorder (ASD) and attentiondeficit/hyperactivity disorder (16). The mutation disrupts the stability of the nucleotide-binding pocket, potentially affecting ATP binding or ADP release (13). The recovery mechanism for such a detrimental mutation in the ATP/ADP-binding pocket remains unclear. Through genetic suppressor screens, biochemical assays, and single-molecule assays, we demonstrate that introducing additional mutations in the nucleotide-binding pocket restores the defects caused by R11Q, likely by restoring the pocket's function. While other substitutions in the motor domain do not restore motility in vitro, they do rescue synaptic and coordination defects in the animal model. Furthermore, treatment with the plant flavonol fisetin improves movement in R11Q worms as well as the in vitro motor activity of the human KIF1A(R11Q) without affecting the wild-type KIF1A, which suggests a potential therapeutic approach for KAND. Our findings shed light on multiple strategies to address R11Q defects at both biochemical and animal levels, offering insights into KAND therapies.

Results

C. elegans Genetic Screens Identified Suppressors for the KIF1A (R11Q) Mutation. To investigate the molecular factors involved in the cellular deficits caused by the disease-associated mutation KIF1A(R11Q), we employed the CRISPR-Cas9 genome editing technique to generate a *C. elegans* model. We introduced the corresponding *unc-104(R9Q)* mutation into the UNC-104::GFP knock-in strain (Fig. 1 *A*–*C*). The resulting *unc-104(R9Q)* mutant animals displayed severe coiling, shortened body length, and pronounced impairments in coordinated movement, with a 100% occurrence rate (Fig. 1*D*, n > 200). These observed phenotypes closely resembled those exhibited by previously reported loss-of-function mutations identified in earlier studies (26).

The complete penetrance of animal defects in the *unc-104(R9Q)* mutant enabled us to conduct a large-scale and high-throughput suppressor screen. However, the unc-104(R9Q) mutant animals exhibited limited mobility and had a reduced brood size compared to wild-type (WT) N2 animals (56.2 \pm 23.8, mean \pm SD, n = 18, approximately one-quarter of the brood size of wild-type animals), which presented challenges in obtaining sufficient synchronized progenies for chemical mutagenesis using conventional methods. To overcome this hurdle, we developed a specialized filter apparatus with a 15-µm diameter that allowed for the enrichment of L1 larvae from mixed population cultures based on their sizes (Materials and Methods). Through this approach, we isolated a total of 103 independent suppressor mutants of unc-104(R9Q) from approximately 100,000 mutagenized haploid genomes. Whole-genome sequencing and bioinformatics analyses revealed 20 intragenic missense mutations within the motor domain, with most of the altered residues highly conserved between C. elegans and humans (Fig. 1 B and C). Additionally, we identified an intergenic suppressor allele, unc-16(E76K) (Fig. 1B), although the rescue efficiency of the intergenic suppressor was weaker compared to the intragenic suppressors (Fig. 1 D and E), it still demonstrated significant effects. Notably, a previously reported missense mutation, L75P, in unc-16 was also found to be a suppressor of other unc-104 partial loss-of-function alleles (27). unc-16 encodes a conserved JNK-signaling scaffold protein regulating directional synaptic vesicle transport, and its dysfunction may cause inappropriate cargo and motor associations which bypass UNC-104 for ectopic synaptic vesicle precursor axonal transportations, probably through kinesin-1 (27).

Through our genetic screens, we identified multiple alleles carrying the same intragenic mutations that exhibited identical rescue effects (Fig. 1B), suggesting that highly recurrent mutations may be responsible for recovering the Unc phenotypes in UNC-104(R9Q). To experimentally validate these identified mutations, we introduced unc-104(T102I) or unc-104(G265R) lesions into wild-type animals using genome editing. Neither of these mutant strains displayed any noticeable uncoordinated behavior, slower swimming speed, or curl morphology (Fig. 1*E*). We then introduced the *unc-104(R9Q)* mutation into these two strains, and interestingly, the resulting double mutant animals did not exhibit Unc defects (SI Appendix, Fig. S1A), which phenocopied the observations made with the unc-104(R9QT102I) and unc-104(R9QG265R) mutants obtained from our genetic suppressor screens. These results confirm that the T102I and G265R amino acid substitutions are indeed intragenic mutations that restore the animal defects caused by the R9Q mutation in UNC-104.

To assess the recovery of worm motility, we conducted an animal swimming assay (25, 26, 30) and compared the paralyzed unc-104(R9Q) animals with the suppressor strains. All suppressors exhibited marked enhancements in animal movement, albeit with varying degrees (Fig. 1E). Among them, we selected two preeminent rescuers, Y53F and T102I, a high-intermediate rescuer A344V, a median rescuer A319T, and a comparatively mild but effective rescuer G265R for subsequent investigations (Fig. 1 F and G and SI Appendix, Fig. S1 B and C). The body length of unc-104(R9Q) mutant animals was only 58% of the WT adults, but the five suppressors restored the body length of unc-104(R9Q) to 72%-94% of the WT level. Notably, T102I and Y53F showed the highest rescue effects, while G265R displayed the least (Fig. 1 D and F). The unc-104(R9Q) mutants exhibited severe coiling and curled postures, with limited spontaneous body elongation (Fig. 1 D and G). We quantified the body curvature using a curling index (SI Appendix, Fig. S1B and Materials and Methods), which was 0.36 ± 0.19 in unc-104(R9Q) mutant comparing to 0.08± 0.03 in WT and found that all five suppressors significantly rescued the curling index, ranging from 0.08 ± 0.05 (Y53F) to 0.10 ± 0.06 (G265R) (Fig. 1 D and G and SI Appendix, Fig. S1C). These results indicate that the intragenic mutations identified within the UNC-104 motor domain effectively rescue the loss-of-function caused by unc-104(R9Q) mutations, and their rescuing effects correlate with animal motility, body length, and morphology.

Biochemical Impacts of Suppressors on the KIF1A Nucleotidebinding Pocket. In order to investigate the effects of the intragenic suppressors on the KAND R11Q mutation in vitro, we examined the motility of purified mouse KIF1A proteins as demonstrated previously (31). The R11Q mutation is located within the nucleotide-binding pocket of the KIF1A motor domain, which is responsible for capturing ATP/ADP molecules (as depicted in Fig. 2A). In the WT motor domain, the arginine (R11) sidechain forms a hydrogen bond with the threonine (T106) sidechain (corresponding to T102 in UNC-104) at the bottom of the nucleotide-binding pocket. These residues wrap around the adenine ring of ATP/ADP, potentially facilitating nucleotide binding (Fig. 2A). Additionally, the tyrosine (Y56) residue (corresponding to Y53 in UNC-104) is located adjacent to R11, and its sidechain forms an additional hydrogen bond with the aspartic acid (D76) sidechain, which likely stabilizes the nucleotide-binding pocket (Fig. 2A).

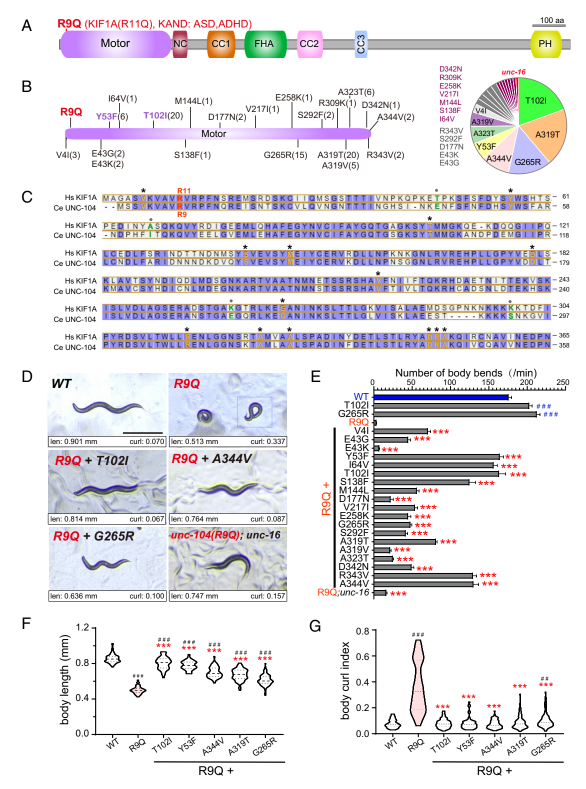
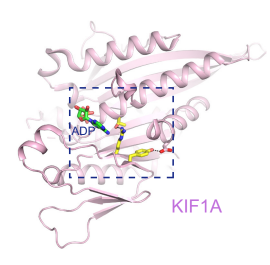
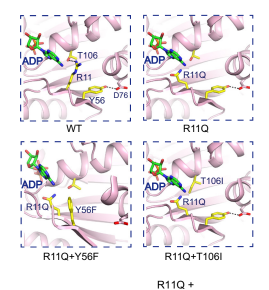
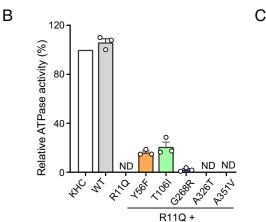
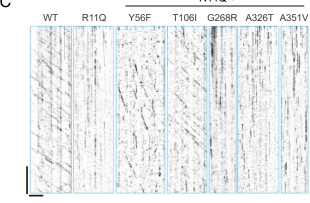


Fig. 1. Genetic suppressor screens for UNC-104(R9Q). (A) Schematic of the domain organization of UNC-104 motor protein. NC, neck coiled-coil. CC, coiledcoil. FHA, forkhead-associated. PH, pleckstrin homology. The R9Q mutation, which corresponds to the KIF1A-associated neuronal disorder (KAND)-associated KIF1A mutation R11Q, was introduced into the UNC-104 protein using the CRISPR-Cas9-based genome editing method. (*B*) *Left*: Amino acid location of isolated R9Q intragenic suppressors within the UNC-104 motor domain. The number of suppressors carrying each mutation site is given in parentheses. *Right*: Pie chart exhibiting the relative population of individual variants among all suppressors. (*C*) Alignment of motor domain sequence between human KIF1A and *C. elegans* UNC-104 proteins. Identical residues are highlighted in purple, and intragenic suppressor sites are labeled with an asterisk or filled circle if they are identical or similar between KIF1A and UNC-104. (*D*) Representative bright field images of WT, *unc-104(R9Q)* mutant, and homozygous suppressor animals at the L4 larval stage. The measured body length (len) and calculated curl index (curl) are shown at the bottom of each image. The data shown in the R9Q image were obtained from the animal framed. (Scale bar, 0.5 mm.) *SI Appendix*, Fig. S1 shows the definition of the curl index and more representative images. (*E*) The number of animal body bends in a water drop was counted for 1 min and shown in columns. Identified UNC-104 amino acid changes in the examined animals are indicated. Error bars represent the SEM, n = 30 animals. Statistical significance was calculated by the unpaired Student's *t*-test: ****P* < 0.001 compared to the animal columoted to the WT animals. (*F* and *G*) Quantification of animal body length (*F*) and curl index (*G*) of five representative suppressors that partially rescued morphological phenotypes of *unc-104(R9Q)* animals. ****P* < 0.001, unpaired Student's *t*-test compared with *unc-104(R9Q)* mutant animals; #









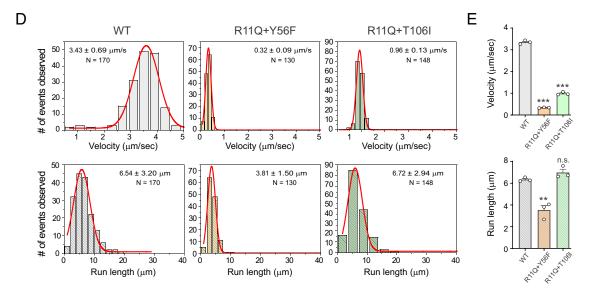


Fig. 2. The structure, ATPase activity assay, and single-molecule study of KIF1A and mutants. (*A*) Potential impacts of mutations on the nucleotide-binding pocket in the motor domain of KIF1A (PDB code: 2ZFI). (*B*) Microtubule-stimulated ATPase activities of the wild-type and various mutants of KIF1A 1-613. Data were normalized by the microtubule-stimulated ATPase activity of kinesin-1 heavy chain (KHC) as 100%. Each experiment was repeated three times independently. Each bar represents the mean value \pm SD, N.D., not detected. (*C*) Representative kymographs showing the motion of wild-type and various mutants of KIF1A 1-613 along the microtubule. Horizontal Scale bar 5 µm; vertical bar 10 s. (*D*) Qualifications of the velocity (*Top*) and run length (*Bottom*) for each protein. The number of events is plotted for the velocity and run length as a histogram and fitted to a Gaussian distribution. The velocity and run length of the corresponding population of the protein (N) are indicated in each panel as mean \pm SD. (*E*) Statistical analyses of the velocity (*Top*) and run length (*Bottom*) for each protein. Statistical significance for the velocity or run length is determined using Student's *t*-test (n.s., not significant; ***P* < 0.001; *****P* < 0.0001) as compared with the WT, n = 3 assays.

However, in the R11Q mutant, the substitution of the longsidechain arginine with a short-sidechain glutamine may disrupt the hydrogen bonding interaction with T106 and hinder the packing with the adenine ring of ATP/ADP (Fig. 2*A*). Consequently, this mutation destabilizes the binding of nucleotides and inactivates the kinesin motor. Consistent with the structural analysis and previous studies (13, 26), the R11Q mutation abolished both the ATPase activity and processive movement of the motor domain of KIF1A (Fig. 2 *B* and *C*).

Among the mutants that restored functionality, the Y56F and T106I substitutions in the nucleotide-binding pocket were able to restore the functionality of the R11Q mutation. In the R11Q and Y56F double mutant KIF1A, through in silico analysis employing PyMOL molecular simulations, we discerned that the Y56F amino acid mutation would disrupt the conventional hydrogen bond between Y56 and D76. Consequently, the side chain of F possibly tends to invert, occupying the spatial cavity induced by the R9Q mutation in the nucleotide-binding pocket of KIF1A (Fig. 2A). This flipping tendency likely compensates for the defect caused by the R11Q mutation. In silico analysis of R11Q and T106I double mutant KIF1A deduced that the substitution of the short-sidechain threonine with a long-sidechain isoleucine in the nucleotide-binding pocket also fills the vacancy created by the R11Q mutation. The T106I substitution forms additional contacts with the adenine ring of ATP/ADP, compensating for the loss of nucleotide binding (Fig. 2A). Therefore, both the Y56F and T106I substitutions in the R11Q mutant KIF1A structurally counteract the mutation-induced defect in the nucleotide-binding pocket.

In our biochemical assays, the Y56F and T106I mutations were found to partially restore both the ATPase activity and processive movement of the motor domain of KIF1A, as indicated by the velocity and processivity measurements (Fig. 2 *B*–*E*). Notably, while the velocity remained compromised, the T106I mutation fully restored the run length of the motor domain (Fig. 2 *D* and *E*), highlighting the positive impact of this substitution in renovating the nucleotide-binding pocket.

It is important to note that three other suppressor mutations, A319T, A344V, and G265R, which are not located in the vicinity of the nucleotide-binding pocket, did not exhibit any significant recovery of KIF1A(R11Q) activity in vitro (Fig. 2 *D* and *E*). This suggests that this group of mutations suppresses the animal Unc defects through mechanisms that are distinct from those of Y56F and T106I, which directly impact the nucleotide-binding pocket.

Restoration of Synaptic Vesicle Distribution and Motility in Suppressors. In our study, we examined the expression pattern and dynamics of UNC-104::GFP in *C. elegans*. In WT animals, we observed UNC-104::GFP fluorescence throughout the nervous system (Fig. 3*A*), which is consistent with previous findings and CeNGEN single-cell RNA-seq datasets (32). However, in animals carrying the UNC-104(R9Q) mutation, we observed a significant reduction in GFP fluorescence intensity, particularly in the nerve ring region (Fig. 3*A*). Although the transcription (*SI Appendix*, Fig. S2*A*) and expression pattern of UNC-104(R9Q) did not show a significant change, the overall fluorescence intensity was diminished.

Remarkably, the intragenic suppressors were able to substantially restore the GFP intensity, indicating their rescuing effects on UNC-104(R9Q) function (Fig. 3*A*). To further investigate the dynamics of UNC-104, we conducted fluorescence recovery after photobleaching (FRAP) experiments in the dorsal nerve cords (Fig. 3*B*, outlined in red). In UNC-104(R9Q) animals, the GFP fluorescence was barely detectable in this region, whereas the genetic suppressors restored GFP fluorescence (*SI Appendix*, Fig. S2 *B* and *C*).

Two of the suppressors, UNC-104(R9QY53F) and UNC-104(R9QT102I), showed similar GFP mobile ratios, indicating the recovery of dynamic behavior comparable to the WT motor (Fig. 3 *B–D*). The GFP recovery ratios for these suppressors were 0.72 ± 0.08 and 0.70 ± 0.10 , respectively, which were in the range of the WT motor (0.68 ± 0.10, mean ± SD) (Fig. 3 *B–D* and *SI Appendix*, Fig. S2*B*). On the other hand, UNC-104(R9QA344V) and UNC-104(R9QG265R), which exhibited partial rescue effects on animal movement (Fig. 1*E*), also partially restored the GFP recovery ratio (Fig. 3 *C* and *D* and *SI Appendix*, Fig. S2*C*). These findings provide further evidence of the rescuing effects of intragenic suppressors on the cellular dynamics of UNC-104 and support their correlation with the partial restoration of animal movement observed in our study.

To investigate the distribution of synapses in different genetic backgrounds, we focused on the DA9 cholinergic motor neuron in the C. elegans dorsal nerve cord (DNC). In WT animals, the DA9 neuron typically forms approximately 25 en-passant synapses along a proximal axon segment in the DNC (SI Appendix, Fig. S3A) (29). We visualized these synapses using a transgenic strain expressing the synaptic vesicle protein (SVP) marker mScarlet::RAB-3 (7, 26, 33) under the control of the DA9 promoter (Pmig-13). In unc-104(R9Q) mutant animals, the DA9 neurons showed a severe deficiency in en-passant synapses (SI Appendix, Fig. S3 A-C). The length of the synaptic region in DA9 neurons was significantly reduced from 115.10 \pm 12.69 μ m in WT animals to 20.48 \pm 26.09 μ m in unc-104(R9Q) animals (SI Appendix, Fig. S3 A and B). Furthermore, the number of mScarlet::RAB-3 puncta, which represents individual synapses, dramatically decreased from an average of 28.36 ± 5.77 in WT animals to only 2.08 ± 2.04 in unc-104(R9Q) animals (SI Appendix, Fig. S3 A and C). Additionally, the size and intensity of mScarlet::RAB-3 puncta were also reduced in unc-104(R9Q) animals (SI Appendix, Fig. S3 D and E). Importantly, the intragenic suppressors were able to significantly restore the distribution area, number, and size of synapses in the DA9 neuron (SI Appendix, Fig. S3 A-E). These findings suggest that the identified suppressor mutations have a rescuing effect on the synaptic defects caused by the *unc-104(R9Q)* mutation.

To investigate whether the axonal synaptic defects in unc-104(R9Q) mutants were attributed to vesicle transport impairments, we examined the motility of mScarlet::RAB-3 puncta in the proximal region of the DA9 axon (SI Appendix, Fig. S3A) (33). By performing kymography analyses, we were able to track and analyze the movement of RAB-3 puncta in both anterograde and retrograde directions. In WT animals, we observed frequent axonal transport of RAB-3 puncta in both anterograde and retrograde directions (SI Appendix, Fig. S4 A, E, and F). However, in unc-104(R9Q) mutants, anterograde RAB-3 transport was disrupted, and there was a significant reduction in retrograde transport events (SI Appendix, Fig. S4 A, E, and F), indicating impaired motility of UNC-104 along the axon. Remarkably, the intragenic suppressors were able to restore the frequency of anterograde and retrograde transport events, as well as the distribution and motility of synaptic vesicles in unc-104(R9Q) mutant animals (SI Appendix, Fig. S4 A-G). This restoration of vesicle transport dynamics explains the recovery of animal Unc phenotypes observed in these suppressors. Notably, the A344V and G265R suppressors, which rescued animal movement but did not restore the motor activity of UNC-104(R9Q) in vitro (Fig. 2 B and C), partially recovered the synaptic defects associated with UNC-104's function in C. elegans (SI Appendix, Figs. S4 A-G and S5F). This suggests that A344V and G265R likely employ distinct rescuing mechanisms from those of Y56F or T106I. Overall,

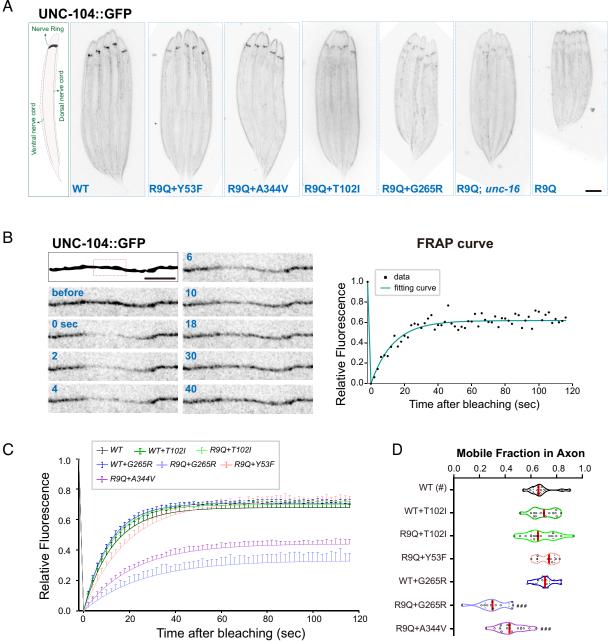


Fig. 3. UNC-104::GFP dynamics in WT and mutants monitored by FRAP. (A) A sketch of the UNC-104::GFP pattern in the wild-type animal, and representative fluorescence images of mutation carrying UNC-104::GFP. GFP was knocked into the C-terminus of UNC-104 before R9Q mutation was introduced, and the unc-104 mutations are indicated in each panel. Note that the fluorescence at the nerve ring was dramatically diminished in the unc-104(R9Q) mutant, and this reduction was rescued to varying degrees in suppressors. (Scale bar, 100 µm.) (B) UNC-104::GFP in axons was photobleached at the boxed area in the top left schematic panel, and recovery was recorded for 2 min. Images before and after photobleaching (at 0 s) at indicated time points are shown in sequential. (Scale bar, 5 µm.) The kinetics of FRAP recovery were fitted with a single exponential equation. (C and D) Quantifications of in vivo UNC-104::GFP FRAP from different variants, showing that suppressors rescued the UNC-104(R9Q)::GFP kinetics (C) and the mobile portion (D) in the axon. Error bars represent SEM, n = 8 to 15 independent animals. Statistical significance was calculated by the unpaired Student's t-test: ###P < 0.001 compared to the WT animals. See also SI Appendix, Fig. S1.

our findings indicate that the intragenic suppressors not only rescue animal motility and morphology, but also restore the synaptic defects and dynamics of synapses in the DA9 cholinergic motor neuron. These results further support the functional significance of the identified suppressor mutations and suggest that targeting some of these sites within the UNC-104/KIF1A motor domain may have potential therapeutic implications for KAND.

Fisetin Alleviates Animal Movement and Morphological Defects in UNC-104(R9Q). Having established the capacity to rescue UNC-104(R9Q) abnormalities through several distinct pathways in vivo, we aimed to explore the potential of small-molecule chemicals in restoring animal behavior. Our accidental starvation of C. elegans carrying the UNC-104(R9Q) mutation led to intriguing observations of improved body morphology and animal movements (SI Appendix, Fig. S6 A and B), which suggests that nutrient conditions may play a role in modulating the behaviors and phenotypes associated with the mutation. This inspired us to explore the potential effects of commercialized food supplements known for their benefits to the nervous system of *unc-104(R9Q)* animals. Our initial focus was on fisetin (34, 35) (SI Appendix, Fig. S6C), a plant flavonol, due to its availability, low cost, and neuroprotective properties. Fisetin has shown effectiveness in improving behavior and physiology in aging mice, preserving survival in C. elegans exposed to thermal stress (36-39). We found that supplementing fisetin in the growth media of C. elegans resulted in improved motility and alleviation of developmental and behavioral defects in *unc-104(R9Q)* mutant animals (Fig. 4A). The effects were dose-dependent, with higher doses showing greater improvements in motility, body length, and posture (Fig. 4 A–D). Though we did not detect a significant restoration of UNC-104(R9Q)::GFP fluorescence in the neurites of animals treated with fisetin (SI Appendix, Fig. S6D), aligning with the modest behavioral amelioration upon fisetin administration, we observed a partial rescue of both the distribution and quantity of neuronal synapses following fisetin treatment (Fig. 4 E and F). This implies that the amelioration of animal behavioral abnormalities may be attributed to the restoration of synaptic functionality. These results underscore the potential of fisetin supplementation as a promising therapeutic strategy for alleviating the symptoms associated with KAND.

cognitive function in Alzheimer's disease models, and promoting

Fisetin Directly Restored the Motor Activity of KIF1A(R11Q)

In Vitro. The observed mitigating effects of fisetin on the unc-104(R9Q) animal prompted us to investigate its potential direct impact on the KIF1A protein. To assess this, we first perform docking analysis between fisetin and the KIF1A motor with the R11Q mutation (Fig. 5A). The docking results revealed the effective binding of fisetin to the adjacent pocket of Q11 in the mutant structure, likely restoring the structural stability of the region surrounding Q11, which was compromised by the R11Q mutation. These findings provide valuable structural insights into the observed functional rescue of the R11Q mutant upon fisetin treatment. Conversely, in the WT KIF1A motor domain, the presence of the R11 guanidine group partially occupied the pocket, thereby hindering the binding of fisetin (Fig. 5 A, Right). These results underscore the specific recognition and functional rescue of KIF1A(R11Q) by fisetin without affecting the wild-type KIF1A motor.

Subsequently, we generated recombinant mouse and human KIF1A(R11Q) proteins (*SI Appendix*, Fig. S7*A*) and conducted biochemical analyses to assess the impact of fisetin on its ATPase activity. We observed negligible ATPase activity in KIF1A(R11Q) treated with 0.1% DMSO, the solvent used for fisetin. However, the presence of 10 nmol/L fisetin led to the detection of ATPase activity of KIF1A(R11Q) to approximately 12% of that in WT KIF1A, the intragenic suppressors Y53F and T102I, which effectively rescued the Unc phenotype caused by R11Q, elevated the ATPase activity of KIF1A(R11Q) to 15 and 22% of the WT level, respectively (Fig. 2*B*). This suggests that even a modest restoration of ATPase activity is potentially sufficient to rescue behavioral defects at the animal level.

Furthermore, we performed a microtubule-gliding assay to examine whether fisetin treatment could induce force generation in KIF1A(R11Q). In the absence of fisetin, KIF1A(R11Q) did not exhibit any MT gliding in vitro. However, the addition of 10 nmol/L fisetin to the gliding assay enabled MT to glide at a speed of 0.49 μ m/s, which is comparable to that of WT KIF1A (*SI Appendix*, Fig. S7*B* and Movie S3). Additionally, using a single-molecule assay, we demonstrated that fisetin treatment enabled KIF1A(R11Q) to generate processive movement along MT, although the speed and run length were lower than those of WT KIF1A (Fig. 5 *C* and *D* and Movies S1 and S2). As predicted by molecular docking, we did not observe any effects of fisetin on the ATPase activity or processive movement of the wild-type KIF1A (*SI Appendix*, Fig. S7*C*). These findings indicate that fisetin specifically enhances the functionality of the KIF1A(R11Q) mutant protein while exerting no effects on the wild-type KIF1A.

Discussion

Our unbiased genetic suppressor screens have successfully identified 20 mutations located within the motor domain of UNC-104 that can rescue the detrimental effects caused by the KANDassociated amino acid substitution. Through comprehensive biochemical and single-molecule studies, we have classified these suppressor mutations into two distinct groups. The first group, represented by Y56F and T106I KIF1A, is situated in the vicinity of the nucleotide-binding pocket. These mutations restore the ATP/ADP binding capacity of KIF1A(R11Q), leading to the recovery of its ATPase activity and motility. In contrast, the second group of suppressors, including A326T, A351V, and G268R KIF1A, does not exert any influence on the nucleotide-binding pocket and fails to restore the biochemical and biophysical properties of KIF1A(R11Q) in vitro. Despite the divergent effects observed in vitro, both sets of amino acid substitutions have demonstrated their efficacy in rescuing the abnormalities in synapse distribution and animal behavior associated with the R11Q mutation.

As delineated by the results in Fig. 3, the effect of the UNC-104(R9Q) mutation is at least partially caused by a protein stability issue, leading to a much lower level of mutant UNC-104 protein. In alignment with this, prior research has posited that UNC-104 protein is susceptible to degradation when its motor functionality was impaired (40). Concurrently, the intragenic suppressors we've identified reside within the ATP motor domain; this suggests that these activity-recovering suppressor motors may mitigate the protein degradation induced by the R9Q mutation. Thus, we suggest that both protein activity and stability may be implicated. These underscore the remarkable potential for restoring function even in the presence of a deleterious KAND mutation that completely abolishes the motor's ATPase activity. Furthermore, although our FRAP and synaptic localization findings (Fig. 3 C and D and SI Appendix, Fig. S5 A-E) did not demonstrate any marked augmentation in motor activity attributable to T102I or G265R, we acknowledge the potential involvement of hitherto unidentified factors that might influence either mutation, leading to increased body bending. The involvement of multiple distinct mechanisms in the restoration process provides great inspiration and suggests that effective interventions for KAND are not only feasible but can be achieved through diverse strategies.

The suppressors belonging to group one suggest that if a secondary mutation can counterbalance the structural defects induced by the primary mutation, it has the potential to restore motor activity. In other words, it may be possible to rescue the defects resulting from the primary mutation even without directly correcting it, as long as a second mutation can counteract its structural effects. For instance, the hydrogen bond normally made by Y56 may not be critical for the motor structure and function, and the loss of the hydrogen bond frees the F residue to flip over to provide spatial support for the ADP binding pocket formation without affecting the whole protein steadiness. Moreover, we observed that mutations Y53F and T102I can restore animal behavior, as well as the UNC-104(R9Q)::GFP signal intensity and their FRAP kinetics within the nervous system, approximating wild-type in vivo profiles (Fig. 3). However, these mutations only partially recover the motor activity in vitro (Fig. 2). Such findings suggest the possibility that specific biophysical attributes of molecular motors, such as individual motor velocity or processivity at their peak capacities, might not be indispensable for overall

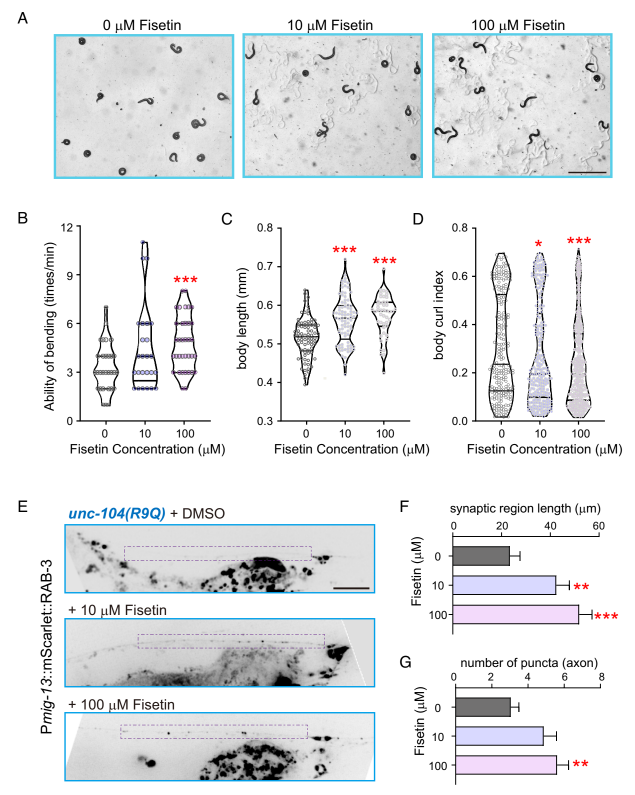


Fig. 4. Fisetin partially recovered the motility and body morphology of *unc-104(R9Q)* animals. (*A*) Representative bright field images of *unc-104(R9Q)* animals treated with different concentrations of Fisetin. (*B–D*). Violin plots harboring raw data dots showing the quantification of animal swimming behavior (*B*, n = 25 to 30 animals), body length (*C*, n = 45 to 85 animals), and curl index (*D*, n =230 to 360 animals) morphology phenotypes of *unc-104(R9Q)* animals treated with different concentrations of Fisetin. Green bars represent the median and quartiles. Statistical significance was calculated by the unpaired Student's *t*-test: *P < 0.05, ***P < 0.001, compared to the *unc-104(R9Q)* animals cultured on the control NGM plates. (*E*) Representative images show the distribution of synaptic vesicles, visualized by P*mig-13*::mScarlet::RAB-3, in the DA9 neuron in *unc-104(R9Q)* mutant with indicated drug treatments. (Scale bar, 10 µm.) (*F* and *G*). Bar graphs showing the quantifications of the length of synaptic region (*P*) and number of puncta (*G*) in the dorsal axon of *unc-104(R9Q)* mutant with indicated drug treatments. *P < 0.01, ***P < 0.001, ***P < 0.001, ***P < 0.001, ***P < 0.001.

organism behaviors and optimal physiological health. Given the pronounced phenotypic revival with motors operating at merely ~ 10 to 50% of the WT metrics, we postulate that a marginal

reinvigoration of motor activity may suffice to restore animal physiology to near wild-type standards. This concept carries significant implications for drug development. A small molecule that mimics

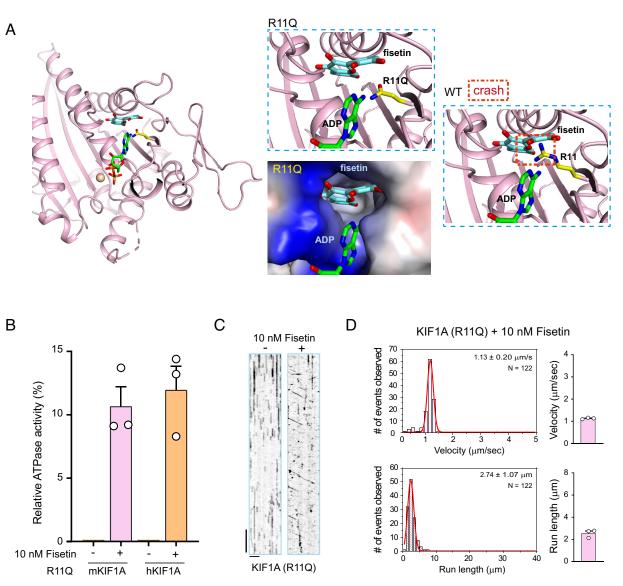


Fig. 5. Fisetin restored the motor activity of KIF1A(R11Q) in vitro. (*A*) Structural model of fisetin docked into the pocket in the KIF1A(R11Q) mutant. *Right* panel, Guanidine group of Arg11 clashes with fisetin. KIF1A PDB code: 2ZFI. (*B*) Microtubule-stimulated ATPase activities of the human and mouse KIF1A 1-613 carrying R11Q mutation, with or without 10 nM fisetin. Data were normalized by the microtubule-stimulated ATPase activity of kinesin-1 heavy chain (KHC) as 100%. Each experiment was repeated three times independently. Each bar represents the mean value \pm SD. (*C*) Representative kymographs showing the motion of mouse KIF1A(R11Q) 1-613, with or without 10 nM fisetin, along the microtubule, respectively. Horizontal scale bar 2 µm; vertical bar 5 s. (*D*) Qualifications of the velocity (*Top*) and run length (*Bottom*) of mouse KIF1A(R11Q) protein with 10 nM fisetin treatment. The number of events is plotted for the velocity and run length as a histogram and fitted to a Gaussian distribution, and the average of three repeats is plotted as a bar graph on the right. The velocity and run length of the protein (N) are indicated in each panel as mean \pm SD.

the effects of the Y56F or T106I substitution in the nucleotidebinding pocket could potentially restore the motor activity of KIF1A(R11Q). Importantly, the T106I single mutation does not result in any detectable abnormalities in *C. elegans*, as shown in Fig. 1*E*. Therefore, such a small molecule-based intervention strategy may not impact the normal function of WT UNC-104/KIF1A but specifically improve the R11Q mutant.

While we have identified several independent mutant alleles of Y53F or T102I, we suspect that our suppressor screens are not yet saturated due to the limitations of the chemical mutagen ethyl methanesulfonate (EMS) used in this study. EMS predominantly induces G/C to A/T nucleotide conversions, thereby only targeting a limited subset of codons (41–43). Future investigations will explore alternative chemical mutagens, such as ENU, to introduce a broader spectrum of mutations for selection. Hence, we speculate that Y56F and T106I are not the sole targets for drug development. As long as the nucleotide-binding pocket can be restored

through the substitution of other residues or by utilizing small molecules, the defects caused by the R11Q mutation can be remedied. Given that mutations in the motor domain of the kinesin family can contribute to various genetic disorders, including those beyond KAND, our design principles hold implications for addressing other kinesin-associated pathologies (1, 3, 44, 45).

Equally intriguing are the suppressors in the second group, which effectively restore synaptic and behavioral defects of *unc-104(R9Q)* in vivo but do not exhibit restored activity of KIF1A(R11Q) in vitro. G268R, as an illustration, showed no improvement of ATPase activity in vitro but strong improvement of vesicle movement in vivo, raising the possibility of whether the rescue of the UNC-104 level is the driving reason for the phenotypic rescue. Our RNA-seq data revealed that the *unc-104* mRNA expression in the R9Q mutant, irrespective of the suppressor presence, remained consistent with the WT levels. This suggests the potential involvement of an elusive regulatory mechanism underlying the G265R or other suppressors-mediated rescue of the R9Q mutation. Considering our current knowledge of motor regulation, it is challenging to conceive a model that can reconcile the differences observed between in vitro and in vivo outcomes. While we have not yet validated all the substitutions in this group as bona fide suppressors, we conducted double knock-in experiments that generated G265R and R9Q double mutant animals de novo, thereby confirming the suppressive effect of G265R on R9Q. The acquisition of multiple alleles through independent screens for many suppressors suggests that we may have identified causative agents capable of restoring R9Q function. These suppressors are likely to act through diverse mechanisms, such as influencing the conformational stability of the protein, modulating the interaction between the motor domain and its cargo, or impacting the recruitment of the motor to the microtubule track. We also propose the existence of certain factors that were not previously recognized for harboring suppressive properties within the second group, which could play a crucial role in activating UNC-104(R9Q), considering the involvement of various intracellular factors that either activate or inhibit motor activity (45-48). One plausible scenario is that an intracellular factor binds to a motif containing G265 and inhibits wild-type UNC-104, and the G-to-R substitution disrupts this interaction, thereby releasing the inhibition in vivo. Alternatively, the G-to-R substitution may lead to the ectopic recruitment of an axonal protein to interact with UNC-104(R9Q), thereby enhancing its motor activity within axons. Future genetic screens involving Unc animals in the background of these rescued animals could potentially shed light on the identification of such molecular candidates. Nonetheless, we assert that motor regulation is likely to be more intricate and fascinating than previously believed, and the exploration of clinical motor mutations will unveil broad avenues for motor research. Our study emphasizes the significance of unbiased genetic

Our study emphasizes the significance of unbiased genetic screens in unraveling the complexities of diseases and devising effective interventions. Although AI has transformed biomedical research by generating numerous hypothetical models, it is noteworthy that AI cannot predict the genetic suppressors identified in this study, despite the availability of KIF1A's structural information. Artificial evolution, on the other hand, which relies on extensive unbiased genetic screens and whole-genome sequencing, holds great promise in complementing AI and providing insights into biology and the understanding of diseases.

The unexpected and inspiring direct effects of fisetin on the UNC-104(R9Q) protein shed light on its therapeutic potential. The kinesin family motor protein has been a subject of drug development efforts over the past decades; however, the efficacy of developed kinesin antagonists or inhibitors has posed limitations for therapeutic interventions, despite their widespread use in laboratories (49). Our findings on fisetin suggest that the development of agonists targeting disease-associated kinesins harboring human genetic mutations could be a fruitful and valuable future direction for motor protein–based drug discovery.

We acknowledge that our study on fisetin in UNC-104(R9Q) animals is still in its preliminary stages. Further experimentation is required to determine the exact interaction between KIF1A and fisetin, and the observed effects of fisetin on improving animal movement or morphology are modest compared to our genetic suppressors. Additionally, fisetin may regulate various complex processes that contribute to the overall physiological improvement in animals. Therefore, we propose considering fisetin as a food supplement rather than a drug for a specific subgroup of KAND patients. Due to its chemical properties and diverse biological effects, fisetin alone may not be an ideal drug; however, the use of this small molecule, along with the proof-of-concept and design principles demonstrated in our study, can provide valuable insights for the development of specific and potent drugs targeting KAND. Considering the reproducible effects of fisetin in both in vivo and in vitro settings, as well as its cost-effectiveness, the application of fisetin as a nutrient supplement may offer potential benefits to KIF1A(R11Q)-associated KAND patients.

Considering fisetin's prospective renovating effects on the nucleotide-binding pocket, it could potentially ameliorate KIF1A variants bearing other mutations within this locus, but not to extraneous domains. From this point of view, precision genomic medicine ought to be refined to the nucleotide resolution: Perturbations at each amino acid within a protein might necessitate unique interventional strategies, underscoring the emerging area of functional residuomics which provides residue-precise functional knowledge of the proteomic landscape.

Caution is crucial when considering the use of fisetin or any other dietary supplement for KAND patients. As mentioned, some KAND mutations have been shown to result in overactive KIF1A (25, 26). Even though our experiments indicated that fisetin exhibited no discernible impact on wild-type KIF1A motor dynamics both in vitro and in vivo, one should be circumspect, as molecules that stimulate the motor might have a dual effect, potentially jeopardizing organism health if they provoke excessive motor activity. Safety considerations, including appropriate dosage, potential interactions with other medications or underlying conditions, should be carefully assessed. We strongly recommend consulting with healthcare professionals or experts in the field of neurodevelopmental disorders before initiating any fisetin supplementation regimen for individuals with KAND.

Materials and Methods

Strains and Genetics. *C. elegans* strains were maintained as described previously (50), on nematode growth medium (NGM) plates with OP50 feeder bacteria at 20 °C. To administer fisetin (Sigma, Cat# 528-48-3), the drug was added to the NGM medium at the desired concentration prior to pouring the plates, which were subsequently seeded with OP50 culture in the usual manner. All the engineered *C. elegans* strains were genetic derivatives of the strain Bristol N2. Strains used in this study are summarized in *SI Appendix*, Table S2. The PHX4787 (UNC-104::GFP) strain, referred to as wild type in this study, and other CRISPR/Cas9-mediated genome editing strains were created by Suny Biotech (Fuzhou City, China). Detailed information such as the modified locus is listed in *SI Appendix*, Table S3. Transformation of *C. elegans* to introduce the *Pmig-13::mScarlet::RAB-3* was performed by DNA injection as described (51), and the information of plasmids and primers is described in *SI Appendix*, Table S4. All animal experiments were performed following governmental and institutional guidelines.

Suppressor Screening. EMS mutagenesis was performed as described before (52, 53) with some modifications. PHX5261 [unc-104(R9Q)::GFP] worms were cultured on regular NGM plates seeded with OP50 feeder till the population was just starved. Then, the mixed-stage worms from 10 to 20 plates were collected, and L1-like larva were sorted with a 15-µm-diameter metal filter. The sorted larvae were cultured to late L4, harvested, and treated with ethyl methanesulfonate (EMS). After the treatment, worms were washed and dispersed to 200-400 OP50-seeded NGM plates by dropping 5 to 10 worms each plate close to the plate margin. Plates were examined when the majority of F2 progenies were older than L4, and basically worms that were able to move to the opposite margin of the plates were very likely to be suppressors. Approximately, 100,000 haploid genomes were checked after multiple runs of screens, and suppressors from different plates were considered independent. After phenotype confirmation over two generations, suppressors were sent to whole genome sequencing followed by comprehensive bioinformation analyses and genetic manipulations to identify and validate the genomic mutations responsible for the rescue.

Imaging and Analysis of the Worm Phenotypes. A scientific cMOS camera (IMX236, Sony Semiconductor Solutions Corporation) was mounted to a compound microscope for bright field images. A swim test was performed as described previously (25, 26, 30) with day 1 young adult worms. Synchronized late L4 stage animals were imaged on NGM plates for body length and curvature analysis. The imaging processing, measurement, and index calculation are described in *SI Appendix*, Fig. S1*B* with ImageJ (http://rsbweb.nih.gov/ij/), and tightly curled animals were not taken into the body length analysis.

Live-Cell Imaging. Synchronized late L4 *C. elegans* animals were mounted on 2% (wt/vol) agarose pads and anesthetized with 0.1 mmol/L levamisole in M9 buffer for live-cell imaging. Our imaging system includes an IX83 inverted microscope (Olympus Lifescience, Inc.) equipped with an EM CCD camera (Andor iXon + DU-897D-C00-#BV-500), and the 488-nm and 561-nm lines of a Sapphire CW CDRH USB Laser System to a spinning disk confocal scan head (Yokogawa CSU-X1 Spinning Disk Unit). The UNC-104::GFP signal of whole animal was imaged with a 10 ×, 0.3NA UPIan Semi Apochromat WD objective. The steady-state images of mScarlet::RAB-3 fluorescent puncta along DA9 neuron axons were taken with $60 \times$, 1.30NA UPIanSApo silicone oil immersion objective at an exposure time of 200 ms, and time-lapse images showing mScarlet::RAB-3 transports were acquired with $100 \times$, 1.49NAApochromat oil immersion objective at an exposure time of 200 ms without interval time for 200 frames. Kymograph extraction, image processing, and measurement of axonal synapse morphology and movement speeds were performed in ImageJ.

FRAP. The fluorescence recovery after photobleaching (FRAP) experiments in *C. elegans* were carried out on the L4 larva worms with GFP reporter knocked into the C-terminus of UNC-104, using a laser scanning confocal microscope (LSM900, Zeiss) equipped with a 63 × ,1.4 NA Plan-Apochromat oil immersion objective. Photobleaching, image acquisition, and raw fluorescence measurement were performed on a ZEN system under identical settings for all experiments. When conducting the assay, the puncta-free GFP-positive dorsal nerve cord processes were photobleached over a rectangular region (65 × 30 pixels, 0.085 µm/ pixel) using 40 iterations of the 488-nm laser with 10% laser power transmission and the pinhole diameter 200 µm. Images were collected with 1% 488-nm laser power over 250 × 60 pixels at 8-bit intensity resolution using a pixel dwell time of 0.96 µs. Prior to the bleaching, two frames of images were acquired, and the recovery of fluorescence was recorded for 120 s with a 2-s interval to guarantee no further recovery was detected. After background subtraction, fluorescence recoveries were calculated as previously described (54).

Protein Expression and Purification. DNA sequences encoding the mouse KIF1A fragment, encompassing the MD-NC-CC1-FHA tandem (residues 1 to 613), were each engineered with a GFP tag adjoined to the C-terminus of the motor protein. These sequences, along with their respective mutants, were subsequently cloned into an adapted pET32a vector, which featured an internal His tag. The human KIF1A protein is derived by mutating the corresponding residues in the mouse KIF1A protein to match their human counterpart, based on the amino acid sequence alignment (SI Appendix, Fig. S7A). All the mutations in the KIF1A fragment were created using the standard PCR-based mutagenesis method and confirmed by DNA sequencing. Recombinant proteins were expressed in Escherichia coli BL21 (codon plus) host cells at 16 °C. After the 16-h cultivation, the cells were harvested and suspended in the buffer containing 50 mM Tris-HCl, pH 7.5, 500 mM (NH4)₂SO₄, 2 mM MgCl₂, 1 mM EGTA, and 1 mM PMSF. The proteins were purified by Ni²⁺-Sepharose 6 Fast Flow (GE healthcare) affinity chromatography with the washing buffer (50 mM Tris-HCl, pH 7.5, 500 mM (NH4)₂SO₄, 2 mM MgCl₂, 1 mM EGTA, 40 mM imidazole) and the elution buffer (50 mM Tris-HCl, pH 7.5, 500 mM (NH4)₂SO₄, 2 mM MgCl₂, 1 mM EGTA, and 500 mM imidazole). The proteins were further purified by size-exclusion chromatography (Superdex-200 26/60, GE healthcare) with the buffer containing 150 mM NaCl, 50 mM Tris-HCl, pH 7.5, 2 mM MgCl₂, 1 mM EGTA, and 1 mM DTT.

Microtubule-Stimulated ATPase Assay. Measurements of the microtubulestimulated ATPase activities of KIF1A MD-NC-CC1-FHA and various mutants were performed by using the HTS Kinesin ATPase Endpoint Assay Biochem Kit (Cytoskeleton, Inc., BK053). Briefly, all of the measurements were based on the malachite green phosphate assay to probe inorganic phosphate generated during the reaction. A standard curve of phosphate was made to estimate the amount of phosphate generated. Each protein sample had three replicates, and each measurement was repeated at least three times independently. The kinesin-1 heavy chain (KHC) supplied in the kit was used as the control. All of the data were analyzed by using the GraphPad prism8 program.

Single-Molecule Assay by Total Internal Reflection Fluorescence Microscopy. Imaging was conducted utilizing a total internal reflection fluorescence (TIRF) microscope (Olympus) equipped with an Andor 897 Ultra EMCCD camera (Andor, Belfast, UK) using a 100 × TIRF objective (NA 1.49, Olympus). The methodologies adopted were described in prior studies (25, 55). In the single-molecule assays, polymerized microtubules were diluted in the taxol-BRB80 buffer and then flowed into a flow cell and incubated for 10 min at room temperature to adsorb onto the coverslip surface coated with anti-tubulin antibodies (Sigma). Subsequently, polymerized microtubules were protected from laser illumination by flowing the basic reaction mix buffer (RM) (0.08 mg/mL glucose oxidase, 0.032 mg/mL catalase, 0.16 mg/mL casein, 1% β-mercaptoethanol, 0.001% tween, 1 mM MgCl₂, 20 µM taxol, and 80 mM D-glucose). Finally, a 50-µL final reaction mix consisting of kinesin motors, RM, and 2 mM ATP was added to the flow chamber, and the time-lapse movies were recorded. Images were taken every 0.1 s with a 0.05-s exposure, with careful modulation of the laser intensity to circumvent photobleaching during consecutive motor runs. For the microtubule gliding assays, purified kinesin motors, diluted 1:40 in RM buffer, were channeled into the flow cell, allowing adherence to the substrate for 5 min. Subsequently, rhodamine-labeled microtubules, diluted 1:40 in a motility mixture (RM complemented with 1 mM ATP), were introduced and visualized using TIRF microscopy.

Single-Molecule Assay Data Analysis. In the single-molecule motility assay, the position of fluorescent motor spots was manually tracked using ImageJ software (Fiji, NIH) (56). In the kymograph, the vertical distance represented the time, and the horizontal distance indicated the run length. The ratio between the run length and the time was the velocity. The number of events for each protein sample was plotted for the velocity and run length as a histogram and fitted to a Gaussian distribution. All the statistical analyses were performed using GraphPad Prism (GraphPad Software), and all the data were presented as mean \pm SD. All the *P* values were calculated by using a two-tailed unpaired Student *t* test.

In Silico Fisetin Docking Analysis. The R11Q mutation was introduced into the KIF1A structure (PDB ID: 2ZFI) using PyMOL (Version 2.5, Schrödinger, LLC). The structures of KIF1A(R11Q) and fisetin were minimized and prepared using BioLuminate software (Maestro BioLuminate 4.4, Schrödinger, LLC, New York, NY, 2021). Induced fit docking was then performed between the two molecules, with the sidechain conformation of Tyr67 and GIn70 allowed to be adjusted.

Data, Materials, and Software Availability. Raw fastq files for RNA-seq from this study have been submitted to the Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo) under accession number GSE252693 (57). All other data are included in the article and/or supporting information.

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